PATENT ABSTRACTS OF JAPAN

(11)Publication number:

54-005888

(43) Date of publication of application: 17.01.1979

(51)Int.CI.

// B05D 5/00

C08F 8/00

(21)Application number: 52-071015

(71)Applicant: MITSUBISHI PETROCHEM CO

LTD

(22) Date of filing:

17.06.1977

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(54) PROCESS FOR PRODUCING HETEROGENEOUS CATION EXCHANGER **MEMBRANE**

(57) Abstract:

PURPOSE: To provide the subject process comprising treating a heterogeneous cation exchange membrane with a resin having an ion-exchange group bridging microcracks produced at the time of after-treatment of said membrane with hot number.

LEGAL STATUS

[Date of request for examination]

[Date of sending the examiner's decision of rejection]

[Kind of final disposal of application other than the examiner's decision of rejection or application converted registration]

[Date of final disposal for application]

[Patent number]

[Date of registration]

[Number of appeal against examiner's decision of rejection]

[Date of requesting appeal against examiner's

decision of rejection]
[Date of extinction of right]

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98本国特許庁

の特許出願公開

公開特許公報

昭54-5888

砂公開 昭和54年(1979)1月17日

MInt. Ci.2		識別記号	
C 98 J	5/22	l	102
B 05 D	5/00		
C 08 F	8/00		

砂日本分類 庁内整理番号
 13(9) F 131 7415-4F
 25(5) K 12 6683-4F
 26(3) F 2 6779-4J

発明の数 1 審査請求 未請求

(全 5 頁)

例不均質カテオン交換膜の製造方法

②特 顧 昭52-71015

顧 昭52(1977)6月17日

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も発明の名称 不均衡カケオン交換膜の製造方法

2.特許糖永の鉱圏

ポリオレフィン樹脂と数粉末状カチオン交換性物質を配合し、得られた混合物を脱状に成形し、この腹状成形物を熱水で処理した後、生じたミタロクラックの一部ないし原表質層にカチオン交換樹脂層を形成させることを特数とする不均質カチオン交換限の製造方法

3.発明の詳細な説明

本発明は改良されたカチオン交換駅の製造方法に関し、特に高イオン農産における時代を対しる時代を対しる時代を対して政力を対して政力を対して政力を対して政力を対して対して対して対している。このようを不均質イオン交換製造自身を製状にした均質イオン交換製造自身を製状にした均質イオン交換製造自身を製状にした均質イオン交換製造自身を製状にした均質イオン交換製造自身を製状にした均質イオン交換製造りも機械的強度が使れている。

がその機械的強度はなか充分ではなく実際に使 用ナるに当つては無々の制剤を免けている。た とえば不均型イオン交換期の矢用性を向上させ るためには、イオン交換機能の化学構造を強固 だするため、果施飯をあめて超麹度を飲くする ことが必要とされているが、イオン交換総密底 が観覚となり必然的に顧の比較抗も増大する。 現去数品化されているイオン交換負は水葱葱に 設治時とこれが空気中で自然乾燥された場合と では彫念収略が激しく影の変形や顔自体にひび が入つたりすることのため実用不能となる。従 つてイオン交換数は限期状態に保つて常温で使 角することが絶対条件となつてなり、そのため **最後として質別する場合の大きを製点となつて** せたとのようなイオン交換胎は一般に乗 映ではなく、従つてこれをイギン交換膜として イオン交換裁獄に使用する場合、四組を除りば かりせなく、イオン交換性を有するという点か **ら興味ある他の用途に使用しようとする場合に** もないがある。

特的写51~ 5888 (2)

とれらの点に解決を与えて到税を不均衡イオ ン交換験を製造することを目的として、マトリ ツクスとしてポリオレフイン街路を用いてとれ と数粉末秋イオン契換物質を混合、成形物、熱 水にて使処理を執す不均質イオン交換膜の楔迹 方法が終案されている。この方法は、例えば特 公附 4 7 - 2 4 2 6 2 号、 特開 图 4 9 - 4 3 8 8 6 号公静、解解的 4 ♀ - 5 3 1 8 ♀ 号明邮祭 等に示されている。しかしたがら上記の方法と より教造した不均質イオン空機期は比較的低い イオン無度の水酔液の脱塩処理用イオン交換膜 としてはもる程度の性値を有しており実用的で はあるが、高いイオン最度の水影響の脱塩処理 用イオン交換組としては性能的にはまだ光分波 足しうるものではなかつた。すなわち、高いイ オン漁鹿の水穏被にないて上記の万法により製 **造した不均質イオン交換膜の比妥抗は十分に像** いがイオン輸車が低下する欠点を有していた。

上記不均質イオン契拠能は無水便処理時代がけるイオン交換関脳の影響に超因してミクロク

ラッタが始生し、とれがイオン交換機の代配の 制御因子の一つとなっているが、このミクロタ ラックが大きいためにこの中に含まれる水また はイオンが高いイオン強能においてイオン強能 を低下させる版因となっていると確定される。

そこで本税明者等は高いイオン機能において 限の比議就を大市化上昇させたいで高いイオン 無準を有する不均質イオン交換筋を得るための 方法を開発すべく福々研究した結果、不均等イオン交換膜を無水で砂湖環する時空じるミクロ オン交換膜を無水で砂湖環する時空じるミクロ 対防で処理することが有効である事実を発見し 本題明に列避した。

すなわち、本発明は高イオン機能における時 イオン輸車の大力る不均偏カチオン交換機を提供することを目的とし、その目的は、ポリオレフイン機能に被粉末状カチオン交換性物質を進合し、待られた協合物を胸状に成形し、この膜状変形物を熱水にて処理した後、生じたミクロクテックの一部ないし段数両層にカチオン交換

数距離を形成らせることを特徴とする高イオン 機成に針ける高イオン軸車の大たる不均変カテ オン交換膜の関連方法により混成することがで きる。

ころにおいて、マトリックス関別として使用されるポリオレフィン関節は、エチレン単独取合体、プロピレン単独取合体、エチレンまたはプロピレン主体の共自合体、かよびこのようなオレフィン単独取合体またな共良合体を主体とする取合体退合物を包含する。

数の末状カテオン交換回旋としては伝激のものが高いられるが、その調製は例えば次のよう にして行なわれる。

(f) 容易にカチオン交換器の導入が可能をビニル 無を有する労務施化合物およびとれと重合能を有する不動和紹合を 2 値以上分子内にほ 有する化合物とを水壁製件中で懸拗失反合させ、 符られたビーズ状央算合体をスルホン化 削て処理し、 とうして得られたビーズ状カチオン交換関節を破扱的に数数許する、

- 回 各品化カリオン交換基の導入が可能及じニル語を有する芳香族化合物およびこれと飲食能を有する不能和総合を2 能以上分子門に保有する化合物とを水供鉄体中で乳化混合させ、物られた砂筋束状炎蛋合体をスルポン化剤で処理して数粉束状カテオン交換機能とする、
- け フェノール化を物ノホルムアルデヒド共和合体を母体としたカチオン交換機能を根据的 に散動弾する、等の方法で行まう。

なから、でいう智易にカチオン交換器の導入が可能なビニル基を有する労務が化合物としては、例えばステレン、ビニルトルエン、ビニルビニルベンセン、α・メテルステレン、ビニルオフタレンまたはその誘連体等の一権または二額以上を用いる。また変合航を有する不動和総合を2個以上分予内に併有する化合物としては、一般に性びビニルベンセンが用いられる。

ポリオレフイン樹脂と飲物取状カチメン交換 樹脂との協合割合は製品の使用目的に応じて定 められるが、一般とは監禁比で2:8~8:2、 このようだして製造された触状の底形物は、60℃以上、好さしくは70℃以上の酸水を用いて欲逸混する。本発的にかける熱水による処理は、比技抗を低下させ、かつ使用中の経験化を少なくする目的でイオン交換機関の影響を可及的大きくするために行なう。したがつて、

(1) カチオン交換性素を有するモノマーと現故 用モノマーかよび場合によつては前記両モノ マーと相称する反応性あるいは非反応性溶鉱 (P) カチオン交換性薬の導入に適したモノマー と架構用モノマー。

カチオン交換性の変としては、カサオも名差であればいずれも適用できる数ママをおければない。カチオン交換にの遊光を有がある。カチオン交換にの変を有が違い、できるなマクリル酸、アクリル酸にアクリル酸にアクリル酸にアクリル酸にアクリル酸にアクリル酸にアクリルをできる。エステル、ビニルスルボン酸エステル等がある。

カテオン交換性素の導入に避したモノマーと してはステレンが一般に使用される。 第合後れ カテオン交換性の遊を導入する方法としては、 公知の方法、例えば健康またはチロル領徴によ るスルホン化労が用いられる。 梅爾斯54-5888 [3]

処理用の熱水は可格性物質を含まないことが弱ましいが、ドクロクラックの発生を制御する必要があるときは、熱水に限、アルカリあるいは 塩化ナトリウム等の質を設加することができる。

一方架機用モノマーとしては、 物えはシビニルベンゼン、 ジメタタリレート 類、 メテレンビスナクリルナミド が出いられる。 カチオン交換性を有するモノマーと 製蔵用モノマーが 組形しない 場合には、アクリル 破ちの反応性のモノマーなよびノまたは水中の神区

応性のモノマーを用いて均一系にして反応を行

て下記の二強りがある。

なり方能が良い。

ればよく、例えはペンソイルパーオキサイド、 ラウロイルパーオキサイド、ライソブルビルパーオキシジカーポオート、ターシャリープテル パーオキシビバレート、アソビスイソプテロニ トリル毎が用いられる。

東合後はモノマーを除去するため化メタノー ルで済を扱、さら化水洗する。 総合化よつては、 その疑惑水にて舟波後処恵する。

以上のようにして役た不均智力チオン交換解 は高イオン難度水解設中において即の比較抗を 形なりにとなく、大巾にイオン輸車が増大する。

以下実施例により本張則をおらに特配に説明する。なお本発明はこれら契約例に限定されるものではなく任意の変更が可能である。契約例中、部とおはすべて重視による。

[比款例-1]

ステレンタ 2 部に対してジビェルベンゼン (純度 5 5 %) 8 部を加え、過酸化ベンソイル等 を密解として診筋重合法により放状共産合体を 特て、これを発性硫酸でスルボン化して頻酸性

[比較例-2]

腰状成形物のを熱水処理する的に(突旋的・1)と同じ条件でイオン交換物能層を形成せしめ、しかる様々5での熱水に30分間浸漉した。このカテオン交換膜の膜厚、イオン熱帯、比 地抗はそれぞれ040m、078、149 Grow であつた。

(與脏例-2)

ステレンスルッン競力りの代りにピニルメルホン微ソーダを使用した似外は「疾的例-1) と両時の万法によつてカチオン交換換を得た。 との腕を95℃の熱水中に50分間型気後の駅 弾み、イオン輸率、比据説はそれぞれ0.39 mm、 0.86、170 0.4mm でもつた。 特別に51-5888(例カテオン交換部別を挟た。この別価性カテオン交換部別を挟た。この別価性カテメンシンは新聞を振りた。この都別の能イオン交換した。この都別のにイオンを整けなるのでは、シートの別がであった。この別がでは、シートの別ができるを対し、シートの別ができるを対し、シートの別ができません。この別が、アニーの別には、アニーの別が、アニーの別が、アニーの別が、アニーの別が、アニーの別が、アニーの別が、アニーの別が、アニーの別が、アニーの別が、アニーの別が、アニーの別が、アニーの別には、アニーの別には、アニーの別には、アニーの別には、アニーの別が、アニーの別には、アニーの別が、アニーの別が、アニーの別には、アニーの別が、アニーのの別が、アニーののでは、アニーののでは、アニーののでは、アニーののでは、アニーののでは、アニーののでは、アニーののでは、アニーののでは、アニーののでは、アニーののでは、アニーののでは、アニー

〔我謝部 - 1]

ステレンスルホン酸カリ 5 1 ガ、 アクリル酸2 4 ガ、ノテレンピスプクリルアミド 2 5 ガ、ベンゾイブンメテルエーテル 2 キノベノマー、 /チかよび水 7 0 ガノモノマーよりなる 放を調製し5 0 でにて均一系としたが、不均負カテオン空機遊回を浸染し、 超気 最 B 本 能 B 郷 級 品 圧 水 優ランプ TRPS 1 - 2 0 (出力 2 頭 / ランプ長 2 5

[突納例-3]

メチレンピスアクリルアミドの代タにエテレンクリコールシメタクリレートを使用した以外は「実施例・1] と同様の方法によつてカサポン交換膜を 物た。この線を 9 5 この結水中に 3 o 分別を受疑の股際 多、イオン戦率、比較抗は それぞれ a 4 0 m、 a 8 s 2 s G A m であつか。

(実施例-4)

メチレンピスアクリルフミド単独の代りにメ チレンピスアクリルフミド 9 %とニチレンダリ コールジメタクリレート 1 6 %を使用した以外 は (実施例 - 1] と同様の方法によつてカチオ ン交換数を待た。との版を 9 5 ℃ の熱水中に 8 0 分間受徴疑の複厚み、イオン輸出、比較鋭は それぞれ B 4 2 m 、 B 9 0 、 2 2 E Q om でもつ た。

(夹焰例 - 5)

スナレン9 2 %、 ジビニルベンセン 6 %、 ベ ンゾインメテルエーテル 2 % / モノマーより 左

特屈刑54-5888:5)

[実施例-4]

スチレンスルホン酸カリミ1%、アクリル酸2 4 %、メチレンピスアクリルアミド29%、通硬酸アンモニウム1%ノセノマー、および水フの%ノベノマーよりなる液中に不均質カチオン交換風回を液体し、航気後 8 0 ℃ で 4 時間車位し、吹質艇を得た。との腰を93℃の輸水中に30分間投資後の膜序み、イオン雑翆、比換抗にそれぞれ0 4 3 mm、 8 B 5 、 1 6 0 Ω·m であつた。

(突 规 例 - 7)

スチレンスルホン数カリ51%、アクリル酸

し、脱気後ナメガラッド電子膨脹熱を行ない改 質験を待た。この態を 9 5 での熱水中化 5 0 分 間を言いの臨厚み、イオン概率、比極抗はそれ ぞれ 0 4 1 mm、 0 8 7 、 5 2 0 G-m であつた。 往1) イオン輪率は脱で 0 5 規定の逆化ナトリウム水 密液とを解離し、膜を介して調水形を膨化等 生する膜電位より弊筋された。

注2) 比級試は Q 5 規定の単化ナトリウムな形式 中において交換電流を適じた時の時の示す電 気銭抗菌 (Q*∞) によつて舞出された。 2 4 %、メテレンピスアクリルブミド2 5 %、および水7 8 % / モノマーよ 5 なる 巻を m 数し 5 0 で に て 片一 果とした 食、 不 海 質 カチ オン 交 後 膜 倒 を 投 資 し、 散 類 は 5 メ メ ラ ツ ド 本 子 領 窓 射 を 行 な い 改 質 級 を 神 た。 と の 摩 を 9 5 ℃ の 駄 水 中 に 3 0 分 間 長 後 長 の 顧 摩 み、 イ オ ン 報 率 、 比 抵抗 は それ ぞれ a 4 2 m 、 a 9 0 、 2 0 0 a ・ m で も つ た 。

[祭舱例-8]

【尖拖狗-9】

アクリル酸 7 5 %、メテレンピスアクリルア
ミド2 5 %、 およびペンジインメテルエーテル
2 第ノセノマーよりなる在を誤製したが、不均
質カチオン交換原因を受強し、 既然後(実施例
- 1 〕と同様の診断にて 2 0 秒限射し改良験を 得た。との膜を 9 5 での際水にて 3 0 分開授減 便の 段降 5 、イオン 報事、 比抵抗は それぞれ 6 は 2 mm、 0 9 2、 3 5 9 Ω·m でもつた。・

がリアクリル世水勘報(ポリアクリル版:水 = 1 : 1) 中に不均型カチョン交換級図を改備

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(19): JAPANESE PATENT OFFICE (JP)

PUBLICATION COUNTRY (10):

DOCUMENT NUMBER (11): Japanese Laid-Open Patent Application

(Kokai) No. 54-5888

DOCUMENT KIND (12): Official Gazette for Laid-Open Patent

Applications (A)

(13):

(15):

APPLICATION NUMBER (21): 52-71015

APPLICATION DATE (22): June 17, 1977

PRIORITY (33) (32) (31):

DISCLOSURE DATE (43): January 17, 1979

NUMBER OF INVENTIONS 1

REQUEST FOR EXAMINATION Not yet submitted

(TOTAL OF 5 PAGES [IN ORIGINAL])

PUBLICATION DATE (44):

PUBLICATION DATE OF NOTICE OF PATENT

ISSUANCE (45):

REGISTRATION DATE (47):

INTERNATIONAL

PUBLICATION NUMBER (87):

ITC (51)²: C 08 J 5/22// B 05 D 5/00 C 08 F 8/00 **CLASSIFICATION SYMBOLS**

102

DOMESTIC CLASSIFICATION (52):

13(9) F 131

25(5) K 12

26(3) K 2

INTERNAL OFFICE

REGISTRATION Nos.

7415-4F

6683-4F

6779-4J

ADDITION TO

(61):

DESIGNATED CONTRACTING

STATES

(84):

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PRIOR ART DOCUMENTS USED IN DETERMINING

PATENTABILITY

(56):

TITLE

(54): Method for Manufacturing
Heterogeneous Cation Exchange
Membrane

ABSTRACT

(57):

SPECIFICATION

1. Title of the Invention

Method for Manufacturing Heterogeneous Cation Exchange Membrane

2. Claims

A method for manufacturing a heterogeneous cation exchange membrane, characterized in that a polyolefin resin is mixed with a finely powdered cation exchangeable substance, the mixture thus obtained is molded into a membrane, and this molded membrane is treated with hot water, after which a cation exchange resin layer is formed on the membrane surface layer or part of the microcracks thus formed.

3. Detailed Description of the Invention

The present invention relates to an improved method for manufacturing a cation exchange membrane, and more particularly relates to a method for manufacturing a heterogeneous cation exchange membrane with a high cation transport number at a high ion concentration. One type of ion exchange membrane comprises an ion exchangeable substance finely dispersed in a synthetic resin matrix in the form of a membrane, and this has been put to practical use as a heterogeneous ion exchange membrane. A heterogeneous ion exchange membrane such as this has better mechanical strength than a heterogeneous ion exchange membrane produced by forming the ion exchange resin itself into a membrane, but the mechanical strength is still not adequate, and various limitations are encountered in actual use. For instance, in order to enhance the practicality of a heterogeneous ion exchange membrane, it is said that the degree of swelling has to be lowered by raising the degree of cross linking so as to strengthen the chemical structure of the ion exchange resin, but this sacrifices ion exchange group density, and the specific resistance of the membrane increases as a matter of course. The ion exchange membranes that are on the market today undergo serious swelling when dipped in an aqueous solution and serious shrinkage when dried naturally in air, so much so that the membrane becomes deformed or cracked and cannot be put to practical use. Therefore, it is an absolute requirement that the ion exchange membrane be stored in a moist state and used at normal temperature, and this poses a major obstacle to use in an apparatus. In addition, since such an ion exchange membrane generally lacks flexibility, not only are there difficulties when using this ion exchange membrane in an ion exchange apparatus, but obstacles are also met in considering use in other applications for which [this ion exchange membrane] would be interesting because of its ion exchangeability.

In an effort to solve these problems and manufacture a novel heterogeneous ion exchange membrane, a method has been proposed for manufacturing a heterogeneous ion exchange membrane by using a polyolefin resin as a matrix, mixing this with a finely powdered ion exchangeable substance, molding this mixture, then subjecting this product to post-treatment with hot water. This method is disclosed, for example, in Japanese Patent Publication 47-24262, Japanese Laid-Open Patent Application 49-43888, and Japanese

Patent Application 49-53189. Nevertheless, while a heterogeneous ion exchange membrane manufactured by the above method was practical in that it did offer a certain amount of performance as an ion exchange membrane for the desalting of aqueous solutions with relatively low ion concentrations, its performance was still lacking when it was used as an ion exchange membrane for the desalting of aqueous solutions with high ion concentrations. Specifically, in an aqueous solution having a high ion concentration, the heterogeneous ion exchange membrane prepared by the above method shows an adequately low specific resistance, but has at the same time a disadvantage that the ion transport number is considerably lowered.

In the above heterogeneous ion exchange membranes, microcracks are formed due to swelling of the ion exchange resin during the post-treatment with hot water, and this is a control factor of the performance of the ion exchange membrane. It is surmised that because these microcracks are fairly large, water or ions contained therein are the cause of the lowered ion transport number at high ion concentrations.

In view of this, the inventors conducted research aimed at developing a method for obtaining a heterogeneous ion exchange membrane that has a high ion transport number without greatly raising the specific resistance of the membrane at a high ion concentration, and as a result they arrived at the present invention upon discovering the fact that it is effective to treat the microcracks that occur during hot water post-treatment of a heterogeneous ion exchange membrane with a resin having crosslinked ion exchangeable groups.

Specifically, it is an object of the present invention to provide a heterogeneous ion exchange membrane with a high cation transport number at a high ion concentration, and this object can be achieved by a method for manufacturing a heterogeneous cation exchange membrane with a high cation transport number at a high ion concentration, characterized in that a polyolefin resin is mixed with a finely powdered cation exchangeable substance, the mixture thus obtained is molded into a membrane, and this molded membrane is treated with hot water, after which a cation exchange resin layer is formed on the membrane surface layer or part of the microcracks thus formed.

Polyolefin resins that can be used as the matrix resin here include ethylene homopolymers, propylene homopolymers, copolymers primarily consisting of ethylene or propylene, and copolymer mixtures primarily consisting of one of these olefin homopolymers or copolymers.

Any finely powdered cation exchange resin can be used, but the preparation thereof is carried out as follows, for example:

- (A) An aromatic compound that has vinyl groups allowing for the easy introduction of cation exchange groups is subjected to suspension copolymerization in an aqueous medium with a compound that has two or more unsaturated bonds per molecule and that can be polymerized with this aromatic compound, the copolymer beads thus obtained are treated with a sulfonation agent, and the cation exchange resin beads obtained in this manner are mechanically pulverized.
- (B) An aromatic compound that has vinyl groups allowing for the easy introduction of cation exchange groups is subjected to emulsion polymerization in an aqueous medium with a compound that has two or more unsaturated bonds per

molecule and that can be polymerized with this aromatic compound, and the finely powdered copolymer thus obtained is treated with a sulfonation agent to obtain a finely powdered cation exchange resin.

(C) A cation exchange resin whose matrix is a copolymer of a phenol compound and formaldehyde is mechanically pulverized.

Examples of the "aromatic compound that has vinyl groups allowing for the easy introduction of cation exchange groups" referred to here include styrene, vinyltoluene, ethylvinylbenzene, α-methylstyrene, vinylnaphthalene, and derivatives of these. These compounds can be used singly or in combination. Divinylbenzene is generally used as the "compound that has two or more unsaturated bonds per molecule and that can be polymerized."

The mixing ratio of the polyolefin resin and the finely powdered cation exchange resin is determined as dictated by the intended use of the finished product, but is generally 2:8 to 8:2 by weight, with a preferable range being 4:6 to 7:3 by weight. The mixing of the polyolefin resin and the finely powdered cation exchange resin can be accomplished by any method that allows the two components to be mixed uniformly, but kneading is generally performed using a roll or an extruder. Particularly desirable is thorough kneading at a temperature over the melting point of the polyolefin resin. Other auxiliary components such as antioxidants, colorants, fillers, and lubricants can be added as needed during this kneading or at any other stage. The obtained mixture is then molded into a film or sheet under suitable conditions in an ordinary roll or press. The plasticizing step that comes before the molding work can be replaced by the step in which the above-mentioned components are kneaded.

The film-shaped article manufactured in this way is subjected to a post-treatment with hot water at 60°C or higher, and preferably 70°C or higher. The hot water treatment is performed in the present invention in order to increase the swelling of the ion exchange resin as much as possible for the purpose of lowering the specific resistance and minimizing changes over time as the product is used. It is therefore preferable for the hot water used in the treatment not to contain any soluble substances, but if it is necessary to control the formation of microcracks, an acid, an alkali, or a salt such as sodium chloride can be added to the hot water.

A cation exchange resin layer is then formed on the membrane surface layer or the microcracks in the ion exchange resin membrane that has undergone this treatment. This cation exchange resin layer can be formed, for example, by a method in which a polymer that has cation exchangeable groups, such as a polymer of acrylic acid, or a polymer that has groups allowing the introduction of cation exchangeable groups is dissolved in a solvent, and [this solution] is then applied to the ion exchange resin membrane by a suitable means such as dipping, after which [this coating] is crosslinked by irradiation, or by a method in which a cation exchangeable monomer and a crosslinking monomer are applied and then polymerized, or a monomer suited to the introduction of cation exchangeable groups and a crosslinking monomer are applied and polymerized, and cation exchange groups are then introduced. The latter method, which starts from monomers, is preferred, however.

Specifically, the combination of monomers can be broadly classified into two groups as follows.

- (A) Monomers having cation exchangeable groups and crosslinking monomers, and in some cases reactive or non-reactive solvents that are miscible with both of the above monomers.
- (B) Monomers suited to the introduction of cation exchangeable groups and crosslinking monomers.

Any groups able to undergo cation exchange can be employed as the cation exchangeable groups, but sulfonic acid groups are preferred for the purpose of preventing an increase in specific resistance. Examples of monomers having cation exchangeable groups include acrylic acid, acrylic salts, acrylic esters, methacrylic acid, methacrylic salts, methacrylic esters, styrenesulfonic acid, styrenesulfonic salts, styrenesulfonic esters, vinylsulfonic acid, vinylsulfonic salts, and vinylsulfonic esters.

Styrene is generally used as the monomer suited to the introduction of cation exchangeable groups. A known method, such as sulfonation by sulfuric acid or chlorosulfuric acid, can be used as the method for introducing the cation exchangeable groups after polymerization.

Meanwhile, divinylbenzene, a dimethacrylate, methylenebisacrylamide, or the like can be used as the crosslinking monomer. If the monomer having cation exchangeable groups and the crosslinking monomer are not miscible, then it is better to conduct the reaction as a uniform system by using acrylic acid, methacrylic acid, or another such reactive monomer and/or water or another such non-reactive monomer.

A substrate membrane is impregnated with the above-mentioned combination of monomers, after which polymerization is conducted, and the polymerization method can involve the use of ultraviolet rays, heat, radiation, or the like. When UV rays are used, however, a known benzoin compound, a benzophenone, a mercaptan, or the like must be used as a photosensitizer, and when heat is used, a peroxide, an azo compound, or the like must be used as a polymerization initiator. In view of the heat resistance of the substrate membrane, that is the heat resistance of the polyolefin, the polymerization temperature should be 100°C or lower, and preferably 80°C or lower. Accordingly, the initiator used when thermal polymerization is conducted should be one that will decompose at this temperature, examples of which include benzoyl peroxide, lauroyl peroxide, diisopropyl peroxydicarbonate, tert-butyl peroxypivalate, and azobisisobutyronitrile.

After polymerization, [the product] is washed with methanol and then with water in order to remove the monomers. In some cases, a post-treatment with hot water is performed once more after this.

A heterogeneous cation exchange membrane obtained in the above manner will have a greatly increased ion transport number without any loss of the specific resistance of the membrane in an aqueous solution with a high ion concentration.

The present invention will now be described in further detail through working examples. The present invention is not limited to these working examples, and modifications can be made within the scope of the present invention. All parts and percentages in the working examples are by weight.

Comparative Example 1

8 parts of divinylbenzene (55% purity) was added to 92 parts of styrene, and copolymer particles were obtained by suspension polymerization using benzoyl peroxide or the like as

a catalyst. This product was sulfonated with fuming sulfuric acid to obtain a strongly acidic cation exchange resin. This strongly acidic cation exchange resin was ground to a grain size of 325 mesh or less in a vibrating ball mill. This resin had a total exchange capacity of 4.5 meq/g on dry base. 40 parts of polypropylene (MI = 10) powder was added to 60 parts of this ground strongly acidic cation exchange resin and subjected to thorough agitation and mixing, after which this mixture was sheet-molded to obtain a molded membrane (A). This molded membrane (A) was soaked for 30 minutes in 95°C hot water, which yielded a heterogeneous cation exchange membrane (B). The thickness of this cation exchange membrane was 0.38 mm, the ion transport number was 0.76, and the specific resistance was $150 \Omega \cdot cm$.

Working Example 1

A solution composed of 51% potassium styrenesulfonate, 24% acrylic acid, 25% methylenebisacrylamide, 2%/monomer benzoin methyl ether, and 70%/monomer water was prepared and made into a homogeneous system at 50°C, after which the heterogeneous cation exchange membrane (B) was immersed in this solution, and after deaeration, this product was irradiated for 10 seconds at an irradiation distance of 10 cm using a Type I-20 high pressure mercury vapor lamp made by JEOL (output: kW, lamp length: 25 cm), which yielded a modified membrane. The thickness of this membrane was 0.45 mm., its ion transport number was 0.94, and its specific resistance was 230 $\Omega \cdot$ cm. This membrane was soaked for 30 minutes in 95°C hot water, after which its thickness was 0.43 mm, its ion transport number was 0.93, and its specific resistance was 180 $\Omega \cdot$ cm.

Comparative Example 2

An ion exchange resin layer was formed under the same conditions as in Working Example 1 prior to the hot water treatment of the molded membrane (A), after which this product was soaked for 30 minutes in 95°C hot water.

The thickness of this cation exchange membrane was 0.40 mm, its ion transport number was 0.78, and its specific resistance was $160 \Omega \cdot \text{cm}$.

Working Example 2

Other than using sodium vinylsulfonate in place of the potassium styrenesulfonate, a cation exchange membrane was obtained by the same method as in Working Example 1. This membrane was soaked for 30 minutes in 95°C hot water, after which its thickness was 0.39 mm, its ion transport number was 0.86, and its specific resistance was 170 $\Omega \cdot$ cm.

Working Example 3

Other than using ethylene glycol dimethacrylate in place of the methylenebisacrylamide, a cation exchange membrane was obtained by the same method as in Working Example 1. This membrane was soaked for 30 minutes in 95°C hot water, after which its thickness was 0.40 mm, its ion transport number was 0.88, and its specific resistance was $230~\Omega$ cm.

Working Example 4

Other than using 9% methylenebisacrylamide and 16% ethylene glycol dimethacrylate in place of methylenebisacrylamide alone, a cation exchange membrane was obtained by the same method as in Working Example 1. This membrane was soaked for 30 minutes in 95°C hot water, after which its thickness was 0.42 mm, its ion transport number was 0.90, and its specific resistance was 220 Ω · cm.

Working Example 5

The heterogeneous cation exchange membrane (B) was immersed in a solution composed of 92% styrene, 8% divinylbenzene, and 2%/monomer benzoin methyl ether, after which this product was irradiated for 20 seconds using the same irradiation apparatus as in Working Example 1. This membrane was sulfonated for 6 hours at 40°C in a solution composed of 83% sulfuric acid (98%) and 17% tetrachloroethane. This membrane was then soaked for 30 minutes in 95°C hot water, after which its thickness was 0.44 mm, its ion transport number was 0.88, and its specific resistance was 230 Ω · cm.

Working Example 6

The heterogeneous cation exchange membrane (B) was immersed in a solution composed of 51% potassium styrenesulfonate, 24% acrylic acid, 25% methylenebisacrylamide, 1%/monomer ammonium persulfate, and 70%/monomer water, and after deaeration, this product was polymerized for 4 hours at 80°C to obtain a modified membrane. This membrane was soaked for 30 minutes in 95°C hot water, after which its thickness was 0.43 mm, its ion transport number was 0.85, and its specific resistance was $160 \Omega \cdot cm$.

Working Example 7

A solution composed of 51% potassium styrenesulfonate, 24% acrylic acid, 25% methylenebisacrylamide, and 70%/monomer water was prepared and made into a homogeneous system at 50°C, after which the heterogeneous cation exchange membrane (B) was immersed in this solution, and after deaeration, this product was irradiated with a one-megarad electron beam to obtain a modified membrane. This membrane was soaked for 30 minutes in 95°C hot water, after which its thickness was 0.42 mm, its ion transport number was 0.90, and its specific resistance was 200 Ω · cm.

Working Example 8

A solution composed of 75% acrylic acid, 25% methylenebisacrylamide, and 2%/monomer benzoin methyl ether was prepared, after which the heterogeneous cation exchange membrane (B) was immersed in this solution, and after deaeration, this product was irradiated for 20 seconds using the same irradiation apparatus as in Working Example 1 to obtain a modified membrane. This membrane was soaked for 30 minutes in 95°C hot water, after which its thickness was 0.42 mm, its ion transport number was 0.92, and its specific resistance was 350 Ω cm.

Working Example 9

The heterogeneous cation exchange membrane (B) was immersed in a polyacrylic acid aqueous solution (polyacrylic acid:water - 1:1), and after deaeration, this product was

irradiated with a one-megarad electron beam to obtain a modified membrane. This membrane was soaked for 30 minutes in 95°C hot water, after which its thickness was 0.41 mm, its ion transport number was 0.87, and its specific resistance was 320 Ω · cm.

Note 1: The ion transport number was calculated from the membrane potential generated between a 0.5 N sodium chloride aqueous solution and a 0.005 N sodium chloride aqueous solution when the two aqueous solutions were separated by the membrane.

Note 2: Specific resistance was calculated from the electrical resistance ($\Omega \cdot cm$) exhibited by the membrane when an alternating current was passed through a 0.5 N sodium chloride aqueous solution.

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